R. Oren · J.S. Sperry · B.E. Ewers · D.E. Pataki N. Phillips · J.P. Megonigal

# Sensitivity of mean canopy stomatal conductance to vapor pressure deficit in a flooded *Taxodium distichum* L. forest: hydraulic and non-hydraulic effects

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Abstract We measured the xylem sap flux in 64-yearold Taxodium distichum (L.) Richard trees growing in a flooded forest using Granier-type sensors to estimate mean canopy stomatal conductance of the stand  $(G_s)$ . Temporal variations in  $G_{\rm S}$  were investigated in relation to variation in vapor pressure deficit (D), photosynthetic photon flux density  $(Q_0)$ , and the transpiration rate per unit of leaf area  $(E_{\rm L})$ , the latter variable serving as a proxy for plant water potential. We found that  $G_{\rm S}$  was only weakly related to  $Q_{\rm o}$  below 500  $\mu{\rm mol}~{\rm m}^{-2}~{\rm s}^{-1}$ ( $r^2=0.29$ ), but unrelated to  $Q_0$  above this value. Above  $Q_0$ =500 µmol m<sup>-2</sup> s<sup>-1</sup> and D=0.6 kPa,  $G_S$  decreased linearly with increasing  $E_{\rm L}$  with a poor fit ( $r^2=0.31$ ), and linearly with  $\ln D$  with a much better fit ( $r^2=0.81$ ). The decrease of  $G_{\rm S}$  with lnD was at a rate predicted based on a simple hydraulic model in which stomata regulate the minimum leaf water potential. Based on the hydraulic model, stomatal sensitivity to D is proportional to stomatal conductance at low D. A hurricane caused an  $\sim 41\%$ reduction in leaf area. This resulted in a 28% increase in  $G_{\rm S}$  at D=1 kPa ( $G_{\rm Sref}$ ), indicating only partial compensation. As predicted, the increase in  $G_{\text{Sref}}$  after the hurricane was accompanied by a similar increase in stomatal sensitivity to D (29%). At night,  $G_{\text{Sref}}$  was ~20% of the daytime value under non-limiting light  $(Q_0 > 500 \ \mu \text{mol} \ \text{m}^{-2} \ \text{s}^{-1})$ . However, stomatal sensitivity to D decreased only to ~46% (both reductions referenced to prehurricane daytime values), thus having more than twice the sensitivity expected based on hydraulic consid-

R. Oren  $(\boxtimes) \cdot B.E.$  Ewers  $\cdot D.E.$  Pataki  $\cdot N.$  Phillips

School of the Environment, Duke University, Durham, NC 27708, USA e-mail: ramoren@duke.edu

Tel.: +1-919-6138032, Fax: +1-919-6848741

R. Oren

The Duke University Wetland Center, Durham, NC 27708, USA J.S. Sperry

Department of Biology, University of Utah, Salt Lake City, UT 84112, USA

J.P. Megonigal

Department of Botany, Duke University, Durham, NC 27708, USA

erations alone. Therefore, non-hydraulic processes must cause heightened nighttime stomatal sensitivity to *D*.

**Keywords** Canopy stomatal conductance · Cypress swamp · Hurricane effect · Light · Nighttime conductance · *Taxodium distichum* · Transpiration · Vapor pressure deficit · Xylem sap flux

## Introduction

The regulation of stomatal conductance  $(g_s)$  by biological and environmental variables has been the subject of intensive research. The relative importance of various control mechanisms is difficult to quantify. In the field, stomatal responses to diurnal changes in photosynthetically active radiation  $(Q_0)$ , vapor pressure deficit (D), leaf temperature, hydraulic conductance within the plant, and soil moisture near the roots are confounded (Sandford and Jarvis 1986). Diurnally, D generally follows the pattern displayed by  $Q_0$ , but with a lag. At the same time, plant and rhizosphere hydraulic conductance tend to decrease, and then slowly increase, mirroring the diurnal patterns of water potential and associated cavitation in the xylem (McCully et al. 1998), and moisture depletion in the soil near roots (Aylmore 1993). In addition, the environment in which plants were grown, and in which measurements are made (e.g., laboratory vs field conditions; Bunce 1998), affects stomatal sensitivity to environmental factors. For example, controlled experiments have been argued to highlight mechanisms of stomatal regulation that may be less important to plants growing in the field (Kramer 1988; Boyer 1989).

Analyzing data collected under moist soil conditions, where soil and soil-root interface resistance are assumed to be small and constant relative to the resistance in other parts of the soil-plant-atmosphere pathway, simplifies analyses of environmental regulation of mean canopy stomatal conductance for water vapor ( $G_S$ ). This allows variation in stomatal conductance to be evaluated solely based on variability in  $Q_o$ , D (Cienciala and Lindroth 1995; Granier and Bréda 1996), and transpiration (Mott and Parkhurst 1991; Monteith 1995), which correlates with bulk plant water potential (Jarvis 1976; Hinckley et al. 1978). However, even under these conditions, resistance to water movement in the rhizosphere increases diurnally in direct relation to the transpiration rate and soil properties, both of which control how fast moisture is depleted near the roots (Kramer 1983; see also Aylmore 1993). This is reflected in the tendency of models to overestimate stomatal conductance in the afternoon (Martin et al. 1997), and the need to use timeof-day as a variable to account for the overestimation (Körner 1993). As a result, the assumption of small and constant root-soil resistance is only partially correct, even under conditions of low transpiration rates, or where plants with high root-to-leaf surface area ratios are monitored while absorbing water from moist, fine soils (Carbon et al. 1980; Sperry et al. 1998). However, under flooded conditions, the assumption of invariable resistance to water uptake from the soil is satisfied.

Flood-tolerant species grow well under flooded conditions, and avoid the adverse consequences of an anaerobic rooting zone mostly by producing adventitious roots into the water above the soil (Sena Gomes and Kozlowski 1980; Pezeshki and Anderson 1997). *Taxodium distichum* (L.) Richard is a highly flood-tolerant species (see Brown and Montz 1986; Megonigal et al. 1997), with transpiration that proceeds at high rates under continuous flooding (Brown 1981; Oren et al. 1999a). Thus, monitoring *T. distichum* under flooded conditions, where there are no diurnal changes in soil resistance to water flow, can supply the necessary data to quantify more accurately stomatal responses to variations in  $Q_0$ , *D*, and transpiration rate.

We investigated the regulation of water use in a 64year-old stand of T. distichum growing on a nearly continuously flooded site. Our primary goal was to quantify the effects of D within the canopy, and  $Q_0$  above the canopy, on mean  $G_{\rm S}$  under conditions of constant, near-zero soil water potential ( $\Psi_s$ ). We hypothesized that T. disti*chum* will show significant reductions in  $G_{\rm S}$  as a direct response to increasing transpiration, and indirect response to increasing ln(D) (see Oren et al. 1999b). During the study, a hurricane reduced the leaf area  $(A_{I})$  in the stand by ~45% (Oren et al. 1999a). Although damage to needle surfaces caused by high wind speed can increase epidermal conductance (i.e., the sum of cuticular and stomatal conductance; van Gardingen et al. 1991), the minimum epidermal conductance of damaged needles is less than a tenth of typical stomatal conductance of T. distichum. This allowed us to test an additional hypothesis that stomata can compensate completely for an abrupt natural defoliation of such magnitude.

Under light-saturating conditions and high D, most plants exhibit a reduction of  $G_{\rm S}$  which limits the rate of water loss and development of potentially damaging low leaf water potential ( $\Psi_{\rm L}$ ). The adaptive advantage of this response may in part be the avoidance of dehydrative damage associated with critically low  $\Psi_{\rm L}$  (Meinzer 1993). For example, several studies have suggested that the absence of stomatal regulation would cause excessive xylem cavitation and failure of water transport (Tyree and Sperry 1988; Sperry et al. 1993; Tyree et al. 1993; Saliendra et al. 1995; Cochard et al. 1996; Lu et al. 1996). This "hydraulic constraint" may be particularly important for species adapted to constant and high  $\Psi_{\rm S}$  (Sperry et al. 1998), such as *T. distichum*.

If one goal of stomatal action under light-saturating conditions is the regulation of  $\Psi_L$ , predictions can be made for the response of  $G_S$  to D and  $A_L$ . The interaction between  $\Psi_L$ ,  $G_S$ , and  $A_L$  in response to D and  $\Psi_S$  can be summarized at steady-state (and infinite boundary layer conductance,  $g_{bl}$ ) by the function (modified from Sperry 1995):

$$\Psi_{\rm L} = \Psi_{\rm S} - (G_{\rm S} \times D) \times \left(\frac{A_{\rm L}}{k}\right) \tag{1}$$

where k is the soil-to-leaf hydraulic conductance, and the ratio  $k/A_{\rm L}$  is the corresponding leaf-specific hydraulic conductance. If we assume constant  $\Psi_{\rm S}$ ,  $\Psi_{\rm L}$ , and k, this relationship predicts that trees with higher  $G_{\rm S}$  at low D will exhibit a proportionally more sensitive stomatal closure with increasing D (see theory in Oren et al. 1999b). It also predicts a perfect compensatory response of  $G_{\rm S}$  in response to reduced  $A_{\rm L}$ , as has been demonstrated in experimental leaf area reduction in a nearby pine stand (Pataki et al. 1998). These predictions are evaluated in light of our observations from T. distichum.

#### **Materials and methods**

The study was conducted at the Lost Shoe Swamp, Duke Forest, Durham, North Carolina  $(35^{\circ}52' \text{ N}, 79^{\circ}59' \text{ W})$ , where mean an-nual temperature is 15.5°C, and mean annual precipitation is 1,140 mm. Seedlings of *T. distichum* were planted in 1933 in an impoundment at an elevation of about 130 m above sea level. Although 1996 was dry relative to the mean, the soil in the study area was continuously flooded to about 0.5 m with slowly flowing water.

In a circular plot of 25 m radius positioned in the middle of the stand, a minor component of Fraxinus pennsilvanica (Marsh.) was present in the sub-canopy [56 trees ha-1 with a basal area at breast height (1.3 m above soil surface) of 4.8 m<sup>2</sup> ha<sup>-1</sup>], with crowns mostly of epicormic branches. Small individuals of T. distichum are healthy and support a normal crown. At age 64, T. distichum in the plot ranged in diameter from 0.18 to 0.56 m, and reached a height of about 27 m. T. distichum was fairly uniformly planted, and has a current density of 722 trees ha-1, with a basal area of 73.6 m<sup>2</sup> ha<sup>-1</sup>. Hydroactive xylem (i.e., sapwood) depth was measured on cores taken from ten trees representing the range in diameter. Sapwood depth varied from 20 to 70 mm (mean=47.5 mm, SE 4.6) but showed only a weak tendency to increase with diameter (P=0.12,  $r^2=0.25$ ). Using the mean sapwood depth from the sampled trees and the diameter of all individuals in the plot, sapwood area was 31.7 m<sup>2</sup> ha<sup>-1</sup>. Leaf area index (LAI; estimated with the Canopy Area Analyzer, LAI2000; Li-Cor, Lincoln, Neb.) was 2.2, thus producing a ratio of sapwood area at breast height to leaf area,  $A_{\rm S}$ : $\hat{A}_{\rm L}$  =0.0014 m<sup>2</sup> m<sup>-2</sup> (Oren et al. 1999a, 1999b).

The study was performed on 22–27 August 1996 when data were collected to evaluate the effect of  $Q_0$  and D on  $G_8$ . On 8–14 September, we resumed measurements to test the hypothesis that the  $G_8$  behavior quantified in August was not altered by a considerable stand-wide defoliation caused by hurricane Fran on 5 September (sustained winds of 35 m s<sup>-1</sup>; Wurman and Winslow 1998).

Measured sap flux ( $J_{\rm S}$ , kg H<sub>2</sub>O m<sup>-2</sup> sapwood s<sup>-1</sup>) in the xylem of trees is increasingly used to estimate  $\bar{G}_{S}$  (Köstner et al. 1992; Granier and Loustau 1994; Granier et al. 1996; Phillips and Oren 1998; Oren et al. 1998a, 1998b), based on the assumption that  $J_{\rm S}$ scaled by  $A_{\rm S}:A_{\rm L}$  is equal to transpiration rate per unit of leaf area  $(E_{\rm L})$ . Thus,  $G_{\rm S}$  for conifers or other species with small leaves (Landsberg 1986; Phillips and Oren 1998; Ewers and Oren 2000) can be calculated using the function (modified from Monteith and Unsworth 1990):

$$G_{\rm s} = \frac{\gamma \times \lambda}{\rho \times c_{\rm p} \times D} \times \frac{J_{\rm s} \times A_{\rm s}}{A_{\rm L}}$$
(2)

where  $G_{\rm S}$  is the canopy stomatal conductance for water vapour (m s<sup>-1</sup>),  $\gamma$  is the psychrometric constant (kPa·K<sup>-1</sup>),  $\lambda$  is the latent heat of vaporization (J·kg<sup>-1</sup>),  $\rho$  is the density of air (kg·m<sup>-3</sup>),  $c_p$  is the specific heat of air at constant pressure  $(J \cdot kg^{-1} K^{-1})$ , and D is the vapor pressure deficit (kPa).  $G_{\rm S}$  can than be converted to molar

units (Pearcy et al. 1989). In Eq. 2,  $\frac{J_s \times A_s}{A_L}$ , equals  $E_L$  only if  $J_S$ 

represents the spatially weighted mean flux (Oren et al. 1998a).

Sap flux density was measured with a Granier-type sensor composed of a heated (constant heat flow) and unheated thermocouple pair connected in opposition to measure temperature difference, as described by Granier (1987). The 20-mm-long sensors were installed in the outer sapwood band of ten trees within 15 m of the plot center at 1.3 m above the soil surface. In five of the trees, a Granier-type sensor was installed in the outer 20 mm of the sapwood. In the other five trees, the modified Granier-Phillips type (Phillips et al. 1996) was used. This sensor measures sap flux separately in the outer 20 mm sapwood and the next 20-mm band of xylem (20-40 mm from the cambium), and uses a single unheated reference thermocouple in the sapwood 10-30 mm from the cambium, after verifying that diurnal radial temperature gradients over these depth ranges were insignificant. In addition, we verified the absence of potential error-causing thermal gradients (Oren et al. 1999a).  $J_{\rm S}$  is calculated from the temperature difference, relative to the maximum difference occurring at times of zero flow. Complete information on sensor installation and flux calculations can be found in Granier (1987) and Phillips et al. (1996).

 $J_{\rm S}$  may be used to estimate  $G_{\rm S}$  only after it is converted to a spatially weighted mean flux. A weighted mean  $J_{\rm S}$  was calculated for each tree using the results of an additional study in this stand which evaluated spatial variation in  $J_{\rm S}$  (Oren et al. 1999a). In that study, (1)  $J_{\rm S}$  of small-diameter individuals (mean diameter=0.24 m) was 0.70 that of trees with diameter 0.30 m or greater, and (2)  $J_{S}$  in the northern third of the tree circumference was 0.64 of that in the other two-thirds. Using these ratios, and the relationships between  $J_{\rm S}$  at sapwood depth of 20–40 mm from the cambium and that in the outer 20 mm of the sapwood (which represent a constant ratio of 0.4),  $J_{\rm S}$  measured in each tree was scaled to total flow by the tree, and divided by the sapwood area of the tree to obtain weighted mean  $J_{\rm S}$ . The flow beyond 40 mm in the sapwood was assumed to equal the flow in the 20- to 40-mm depth. Once the mean flux in each tree was obtained, a population mean  $J_{\rm S}$  at half-hourly temporal scale was calculated, and multiplied by the population mean of  $A_S:A_L$  to estimate  $E_L$ , and  $G_S$ (Eq. 2), at the canopy scale.

To calculate D (kPa), air relative humidity and temperature were measured (HMP 35C probe; Vaisala, Finland) at 20 m aboveground, corresponding to 0.67 of the canopy length from the bottom. In addition, before the hurricane,  $Q_0$  (µmol m<sup>-2</sup> s<sup>-1</sup>) was measured above the canopy (LI-190S; Li-Cor) by attaching the sensor to the end of a pole, and extending the telescoping pole above the canopy. Light measurements were unavailable after the hurricane due to sensor failure. The xylem flux and micro-meteorological data were sampled with a multi-channel data logger (Delta-T Devices, Cambridge, UK) at 30-s intervals, averaged and recorded every 30 min.

Three important assumptions in using Eq. 2 for calculating conductance are: (1)  $E_{\rm L}$  is directly related to  $J_{\rm S}$  measured at the lower stem, i.e., the stem hydraulic capacitance is negligible (Jarvis 1976; Phillips and Oren 1998); (2) leaf boundary layer



Fig. 1 Diurnal relationships between ensemble mean sap flux density  $(J_S)$ , weighted for the entire sapwood area, and vapor pressure deficit (D) in the pre- and post-hurricane period (A), and photosynthetic photon flux density  $(Q_0)$  in the pre-hurricane period **(B)**. *Vertical lines* represent 1 SE of the mean (n=10 trees) of the ensemble means. Arrow represents the direction of the hysteresis

conductance  $(g_{bl})$  is large relative to leaf stomatal conductance  $(g_{\rm S};$  Whitehead and Jarvis 1981), and (3) D is uniform throughout the canopy volume, requiring only one measurement point.

In interpreting the diurnal relationship of  $J_{\rm S}$  and D, a counterclockwise hysteresis indicates hydraulic capacitance (Doley 1967; Schulze et al. 1985; Meinzer et al. 1997), in proportion to the width of the hysteresis (Jarvis 1976; Hinckley et al. 1978). The absence of such a hysteresis (Fig. 1A) provides strong support for the first assumption, justifying the use of  $J_{\rm S}$  for estimating  $G_{\rm S}$ . The counter-clockwise hysteresis seen in the relationship between weighted  $J_{\rm S}$  and  $Q_{\rm o}$  (Fig. 1B) indicates that although stomata may open with light, transpiration, as reflected in xylem flux rate, is too low to measure as long as D is very low. Thus, assumption 1 is met in this study.

Assumption 2 is generally considered true for narrow leaves (Whitehead and Jarvis 1981; Landsberg 1986). This assumption is reflected in Eq. 2, where the energy term is neglected, because D is considered approximately equal to leaf-to-air vapor pressure difference, and is calculated from measurements within the canopy volume but outside the leaf boundary layer. Boundary layer conductance,  $g_{b1}$  (mmol m<sup>-2</sup> s<sup>-1</sup> at 25°C), can be estimated using Jones (1992):

$$g_{\rm bl} = 304.4 \times \sqrt{\frac{u}{d_{\rm L}}} \tag{3}$$

where u is mean windspeed (m s<sup>-1</sup>), and  $d_{\rm L}$  is a characteristic leaf dimension (m). Based on the dimension of either a single needle-leaf of *T. distichum* ( $d_L < 0.002$  m) or of the lateral branchlets ( $d_L < 0.03$  m; Brown and Montz 1986), it is possible to solve for the windspeed that will result in  $g_{bl}/g_s=10$ , a condition in which the conductance calculated from  $J_s$  is dominated by the stomata rather than the boundary layer (Whitehead and Jarvis 1981). Using a typical stomatal conductance of T. distichum seedlings (ca 100 mmol m<sup>-2</sup> s<sup>-1</sup>; Pezeshki and Anderson 1997), the windspeed necessary to create this condition is 0.02 and 0.32 m s<sup>-1</sup> for the leaf and branchlet, respectively. In our region, such rates prevail (>90% of the daytime shows windspeed >0.4 m s<sup>-1</sup>, the lowest resolution of a typical cup anemometer). Furthermore, both rates are probably somewhat overestimated because seedlings have higher stomatal conductance than mature, taller individuals of the same species (Friend 1993; Saliendra et al. 1995), and Eq. 3 tends to underestimate  $g_{bl}$  (Grace 1989). Thus, in this stand, high  $g_{\rm bl}$  predominates because it can be attained with low windspeeds, and because wind can penetrate with little obstruction into the relatively open stand.

Although *D* must vary somewhat with height because  $D\approx0$  near the water surface, given the low LAI and the prevailing wind conditions, the air within the canopy can be considered well mixed with the air above the canopy (Jarvis 1976; Parker 1995). This supports the third assumption that *D* measured at one point, at 2/3 canopy height, represents *D* in the entire canopy zone (6–27 m aboveground; e.g., Sullivan et al. 1996).

All statistical analyses were performed using SAS software (SAS Institute, Cary, N.C.).

## **Results and discussion**

Ensemble means for the pre- and post-hurricane periods show that the rate of water uptake at half-hourly resolution increased linearly with D before the hurricane, but followed a lower trajectory with a tendency to saturate in the post-hurricane period (Fig. 1A). Although the relationship between weighted  $J_{\rm S}$  and D did not include a hysteresis commonly seen when lag occurs between Ddriven transpiration and water uptake, a large hysteresis was seen with  $Q_0$  in the pre-hurricane period (Fig. 1B). Thus, uptake begins about 0.5-1.5 h after dawn, when D begins to increase, and continues into the night, until D reaches zero. Variation among trees in lags between  $J_{\rm S}$ and  $Q_0$  were unrelated to tree size (P>0.05). Furthermore, unlike nearby *Pinus taeda*, which showed stomatal closure at night thus using water uptake for recharge (Phillips et al. 1996), water taken up at night by T. disti*chum* is immediately transpired. This behavior was especially clear on 13 September when nighttime flux, mostly in the outer sapwood, accompanied a nighttime surge in D to 0.6 kPa (Oren et al. 1999a). Thus, use and replenishment of water stored in this species is not large, allowing  $G_{\rm S}$  to be calculated from  $J_{\rm S}$  on half-hourly time scales without accounting for modulation by storage. Half-hourly means of  $G_{\rm S}$  under sufficient light and low D were within the range of instantaneous  $g_{s}$  reported for seedlings (Pezeshki et al. 1995; Pezeshki and Anderson 1997). Nevertheless, caution must be used when interpreting early morning data, when  $J_{\rm S}$  and D are often immeasurable, but  $G_{\rm S}$  may be high.

### Daytime responses of $G_{\rm S}$ to $E_{\rm L}$ , $Q_{\rm o}$ , and D

To address the primary objective of this study, it was necessary to relate  $G_S$  to  $E_L$ ,  $Q_o$ , and D. Empirical relationships between  $g_S$  and  $Q_o$  or D provide a convenient approach to describing the response of stomatal conductance to varying atmospheric conditions (Jarvis 1976; Sandford and Jarvis 1986; Whitehead 1998). An exponential rise to a maximum is the form commonly used to describe stomatal response to light (Jarvis 1976). Many forms of non-linear functions have been used to describe the decrease in  $G_S$  with increases in D (see Lohammar et al. 1980; Massman and Kaufmann 1991; McNaughton and Jarvis 1991; Granier and Loustau 1994; McCaughey and Iacobelli 1994; Monteith 1995; Dang et al. 1997; Martin et al. 1997). Models use these functions to generate a multiplier, one of several used in multipleconstraint functions, to reduce maximum  $g_S$  or  $G_S$  to estimates of actual conductance (see Schulze et al. 1994); maximum conductance is determined for each species under non-limiting environmental conditions (e.g., high irradiance without water limitation; Dai et al. 1992).

In analyzing the effect of environmental variables on  $G_S$ , we used an empirical non-linear multiple regression model (Bréda et al. 1993; Granier and Bréda 1996) after Lohammar et al. (1980) to quantify the effects of D and  $Q_0$  on  $G_S$ . The model is modified from the original by replacing global radiation with  $Q_0$ :

$$G_{\rm S} = \left[\frac{Q_0}{Q_0 + a}\right] \times \left[b + (m \times \ln D)\right] \tag{4}$$

where *a*, *b*, and *m* are empirically fitted coefficients.

This empirical model performs as well as biophysically based functions (Massmann and Kaufmann 1991), and may have a better theoretical justification than certain of the other empirical functions in that (1) it is consistent with the assumption that  $g_S$  decreases linearly with  $E_L$  (Leuning 1995; Monteith 1995), indicating a mechanism designed to minimize although not eliminate changes in bulk leaf water potential (Sandford and Jarvis 1986), and (2) it accounts for the effect of the initial maximal value of  $g_S$  on the sensitivity of  $g_S$  to D(McNaughton and Jarvis 1991). The relationship between  $G_S$  and lnD provides the coefficient b, which can serve as a reference  $G_S$  ( $G_{Sref}$  at D=1 kPa), which is especially convenient because it falls within the range of typically available data, thus not requiring extrapolation.

The half-hourly values of  $G_{\rm S}$  were analyzed based on Eq. 4. The model accounted well ( $r^2$ =0.77, P<0.0001) for variation in pre-hurricane  $G_{\rm S}$  (while  $Q_{\rm o}$  was available and before  $A_{\rm S}$ : $A_{\rm L}$  increased), yielding the coefficient estimates a=121.3 (SE 35.2), b=70.6 (SE14.6), and m=35.5 (SE 11.0).

We partitioned the data into two parts in which interactive effects of increasing light and vapor pressure deficit on  $G_{\rm S}$  are minimized in order to test the hypothesized effects of  $E_{\rm L}$  and D on  $G_{\rm S}$ . Transpiration of a T. distichum sapling was unrelated to  $Q_0$  above 200 µmol m<sup>-2</sup> s<sup>-1</sup> (Steinberg et al. 1990). Given the low LAI in this, as in many other T. distichum stands (Brown 1981), light at the bottom of the canopy is about 260  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup> when  $Q_0$ =500  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup>. After selecting data representing  $Q_0 > 500 \ \mu \text{mol m}^{-1} \text{ s}^{-1}$ , which effectively removed data for D < 0.6 kPa, the model (Eq. 4) results were significantly improved ( $r^2=0.86$ , P<0.0001). For the selected data,  $G_{\rm S}$  decreased significantly with increasing  $E_{\rm L}$  (P<0.001; Fig. 2A, open symbols), but the response was significantly better correlated to increasing  $\ln D$  (P<0.0001; Fig. 2B, open symbols). An approach to remove the effect of auto-correlation between  $G_{\rm S}$ ,  $E_{\rm L}$ , and D, proposed by Monteith (1995), yielded the same outcome.  $G_{\rm S}$  for the rest of the data (i.e.,  $Q_{\rm o}$ <500 µmol m<sup>-1</sup> s<sup>-1</sup> or D<0.6 kPa) was weakly related to  $Q_0$ (P<0.005; Fig. 2C), but not to  $E_{\rm L}$  or D.



Based on these results, in the absence of limits to water availability, variability in  $G_{\rm S}$  of T. distichum is mostly related to D at  $Q_0 > 500 \text{ }\mu\text{mol }m^{-2} \text{ s}^{-1}$ , a value of  $Q_0$  that may reflect non-limiting light throughout the canopy. At  $Q_0 < 500 \ \mu\text{mol m}^{-1} \text{ s}^{-1}$ , a more detailed approach may be necessary to describe the variability in  $G_{\rm S}$ , e.g., one that estimates the weighted mean light as a function of leaf area density distribution and light attenuation (Campbell and Norman 1998).

#### Response to defoliation

Hurricanes in T. distichum stands tend to strip foliage and branches (Putz and Sharitz 1991), causing an abrupt reduction in  $A_{\rm L}$ . The reduction in  $A_{\rm L}$  in this stand by hurricane Fran was estimated at ~45% (Oren et al. 1999a). We confirmed this estimate by using conductance values obtained at night. At very low D at nighttime (e.g., 0.14 kPa), stomata were probably as open after the hurricane as before it. Thus, the difference in  $G_{\rm S}$  (Fig. 3A) between the two nighttime data sets at D=0.14 kPa (48.6 vs 28.7 mmol m<sup>-2</sup> s<sup>-1</sup> for pre- and post-hurricane, respectively) is due to not accounting for leaf loss in the *calculations* of post-hurricane  $G_{\rm S}$ . To match the post-hurricane  $G_{\rm S}$  with the pre-hurricane value required a 41% reduction in  $A_{\rm L}$  to reflect leaf loss ( $A_{\rm S}:A_{\rm L}$  increase from 0.0014 to 0.0024 m<sup>2</sup> m<sup>-2</sup>), remarkably similar to the estimate based on leaf fall (45%).

A 41% reduction in  $A_{\rm L}$  will cause an inversely proportional increase (69%) in the leaf-specific hydraulic conductivity  $(k/A_{\rm L})$  assuming no change in k. If we further assume a constant  $\Psi_L$ , all else being equal, Eq. 1 predicts that a 69% increase in  $k/A_L$  will cause (1) a 69% increase in  $G_{\rm S}$ . Furthermore, we have recently shown that under a given set of environmental conditions, stomatal conductance sensitivity to D, quantified as  $dG_S/dlnD$ , increases proportionally to stomatal conduc-



**Fig. 3A–C** Mean canopy stomatal conductance  $(G_S)$  in relation to the natural logarithm of vapor pressure deficit (D). A Nighttime values for the post-hurricane period are shown uncorrected for leaf loss as symbols, and a least-square fit to the corrected data is shown as a dashed line that matches the limited nighttime data in the pre-hurricane period. **B** Daytime response of  $G_{\rm S}$  to D as in **A**, selected for non-limiting light conditions based on a boundary line analysis (see text). Post-hurricane values are corrected for leaf loss (see text). Vertical lines represent 1 SE (n=10)

tance at low D, e.g., at 0.6  $G_{\text{Sref}}$  (Oren et al. 1999b). Thus, if  $G_{\text{Sref}}$  under non-limiting light increases by 69%, (2) stomatal response to  $\ln D$  is predicted to also increase by 69%. Finally, (3) stand transpiration would be the same before and after the hurricane. Such a response was



**Fig. 4** A The response of mean canopy stomatal conductance  $(G_S)$  to vapor pressure deficit (D) during nighttime and during pre- and post-hurricane daytime hours. The *symbols* represent the least-square fit to the data in Fig. 3A, B, and the lines represent a stomatal response predicted based on a simple hydraulic model and  $G_S$  at D=1 kPa ( $=G_{Sref}$ ) (see text). **B** The sensitivity of  $G_S$  to increasing D as a function of  $G_{Sref}$  representing the slope and intercept of the relationship  $G_S=(m \times \ln D)+b$ , respectively. The *diagonal line* is the theoretical relationship between the two parameters, and *symbols* are as in **A**, where a *filled symbol* is a nighttime value (1 SE for both coefficients is shown for each datum; in some cases it is smaller than the symbol)

indeed seen for a neighboring *P. taeda* L. stand over a 45% reduction in  $A_L$  (Pataki et al. 1998). These are the predictions for a perfect compensatory response of  $G_S$  to partial defoliation.

To facilitate a comparison of  $G_{\rm S}$  response to D between the pre- and post-hurricane period, an evaluation of all predictions was made on data selected from scatterplots of  $G_{\rm S}$  versus D, based on a boundary line analysis (Martin et al. 1997; Schäfer et al. 2000). This was necessary because light measurements were not available for a conditional sampling of data in the post-hurricane period. There was no difference in the pre-hurricane period between the relationships obtained using  $Q_{\rm o}$ >500 µmol m<sup>-1</sup> s<sup>-1</sup> and the boundary line analysis, reflecting the lack of sensitivity of *T. distichum* to light above the selected threshold. In addition, post-hurricane  $G_{\rm S}$  was calculated based on a 41% leaf loss.

Both  $G_{\text{Sref}}$  and the stomatal sensitivity to D (d $G_{\text{S}}$ /dlnD) increased following the hurricane (Fig. 3B;  $G_{\text{Sref}}$ =69.0 and 88.4 mmol m<sup>-2</sup> s<sup>-1</sup> for pre- and post-

hurricane, respectively, paired Student *t*-test P=0.03; dG<sub>S</sub>/dlnD=-38.9 and -50.3 mmol m<sup>-2</sup> s<sup>-1</sup> kPa<sup>-1</sup> for preand post-hurricane, respectively, P=0.02). As predicted (2), the increase in both parameters was approximately proportional (28 and 29% increase in G<sub>Sref</sub> and dG<sub>S</sub>/dlnD, respectively). The least-square fit to the data in Fig. 3B, shown as symbols in Fig. 4A (note that D in Fig. 4A is not transformed), matched closely the predictions based on the hydraulic constraint whereby dG<sub>S</sub>/dlnD≈-0.60×G<sub>Sref</sub> (Oren et al. 1999b), shown as lines in Fig. 4A. Thus, the consequence of defoliation by the hurricane was to increase both the conductance at low D and stomatal sensitivity to D along a theoretical line representing stomatal regulation of leaf water potential (Fig. 4B).

The ~30% increase in  $G_{\text{Sref}}$  and  $dG_{\text{S}}/d\ln D$  fell short of the 69% increase predicted for a perfect compensatory response (1). Thus, contrary to prediction 3, stand transpiration decreased by 18% rather than staying constant (Oren et al. 1999a). The reduction in stand transpiration is close to the 24% reduction predicted from the 41% loss of leaf area, compensated by an only 28% increase in  $G_{\text{Sref}}$ , which, according to Eq. 1 (for constant k and  $\Psi_{\rm L}$ ) and Fig. 4A, B, would apply to  $G_{\rm S}$  at all D. Possible reasons that, singly or in combination, could account for the lack of a perfect compensatory response by T. disti*chum* stomata include (1) a 24% reduction in either k or  $\Delta \Psi$  caused by the hurricane, and (2) a maximum possible  $G_{\rm S}$  set by stomatal density and maximum aperture that is lower than the increase in  $G_{\text{Sref}}$  required to compensate for the loss of foliage.

#### Nighttime responses of $G_{\rm S}$ to D

Some species keep their stomata open at night, and transpire in response to D (Green et al. 1989; Becker 1998; Benyon 1999). In species that keep stomata open at night, nighttime sensitivity of  $G_{\rm S}$  to D can first be predicted based on hydraulic considerations alone (Eq. 1, and Oren et al. 1999b). Then, using a departure from the predicted response, non-hydraulic regulation of  $G_{\rm S}$  sensitivity to D can be inferred. In a stand of Salix viminalis L.,  $G_S$  at nighttime was in fact nearly equal to  $G_S$  at daytime (global radiation >400 W m<sup>-2</sup>) when compared at D=0.1 kPa (calculated from Iritz and Lindroth 1994). However, at D=0.5 kPa, nighttime  $G_S$  was only 15% of daytime values thus showing a disproportionate increase in nighttime sensitivity to D. A similar increase in sensitivity to D at night was found in a stand of Populus tremuloides Michx. (Hogg and Hurdle 1997). These findings indicate that stomatal sensitivity to D at night increases more than would be predicted based on hydraulic considerations regarding the role of stomata in regulating leaf water potential.

Nighttime transpiration in a solitary sapling of T. distichum has been shown to have a similar pattern as air temperature, and thus D (Steinberg et al. 1990). We therefore searched the nighttime record for data that could be used to evaluate whether nighttime stomatal sensitivity to D is higher than can be expected based simply on hydraulic constraints.

We extrapolated daytime  $G_{\rm S}$  to D=0.14 kPa under non-limiting light, using the pre-hurricane relationship (Fig. 3B), and compared to the night values obtained during the pre-hurricane period at the same D (Fig. 3A).  $G_{\rm S}$  of T. distichum at night averaged about 40% of that in the daytime at D=0.14 kPa (Student-*t* P<0.01). Cuticular conductance in several conifers (Hadley and Smith 1990) was at least three orders of magnitude smaller than the nighttime  $G_{\rm S}$  found here, and does not respond to D, and so could not corrupt the analysis of nighttime  $G_{\rm S}$ . Because  $G_{\rm S}$  at a given D is lower at night, based on Eq. 1, stomatal regulation of  $E_{\rm L}$  with increasing D does not need to be as strict to prevent  $\Psi_{\rm L}$  from reaching damaging (e.g., cavitation-inducing) values (Oren et al. 1999b). Thus, based on hydraulic considerations, we predicted that  $G_{\rm S}$  at nighttime would show 40% of the daytime sensitivity to D. This prediction was evaluated with post-hurricane data (full line in Fig. 3A) corrected for leaf loss (dashed line in Fig. 3A).

Comparing the nighttime  $G_{\rm S}$  response to D [ $G_{\rm S}$ =  $13.4-(17.8\times\ln D)$ ; dashed line in Fig. 3A] with the prehurricane daytime response [ $G_S$ =69.0–(38.9×lnD); circles in Fig. 3B] shows that the hydraulically based proportionality (~0.6; Oren et al. 1999b) between  $dG_S/dlnD$ (the slope in the equations above) and  $G_{\text{Sref}}$  (the intercept) is kept during the daytime (=0.56) but is far exceeded at night (=1.33). This is clearly demonstrated in that the least-square fit to the nighttime data (depicted as full circles in Fig. 4A) shows a greater stomatal sensitivity to D than is predicted by the hydraulic model (the respective line in Fig. 4A), which uses  $0.6 \times G_{\text{Sref}}$  as a proportionality for the D sensitivity of  $G_{\rm S}$ . This contrasts with the good correspondence between the least-square fit and model predictions during daytime both before and after the hurricane (Fig. 4A). Thus, the nighttime  $G_S$  of T. distichum decreased from 40% of daytime values at D=0.14 kPa to about 30% at D=0.5 kPa, and is projected to reach zero at D=2.1 kPa. This stomatal behavior of T. distichum shows a sensitivity to D at nighttime that is twice the sensitivity predicted based on hydraulic considerations, as is demonstrated by the position of the nighttime  $G_{\rm S}$  datum above the theoretical hydraulically controlled response (full circle in Fig. 4B; P<0.001).

The three species, *T. distichum*, *S. viminalis* and *P. tremuloides* function and grow well in moist soils, and keeping stomata open when soil moisture is not limiting may simply reflect the absence of a need to conserve moisture. When light becomes available in the morning, these mesic and hydric species may, more than other species, avoid the commonly observed lag between enhanced assimilation and increased stomatal conductance (Körner 1993; Pearcy et al. 1994). The tendency of *T. distichum* to form stands of low LAI, thereby having stomata reaching maximum conductance at low  $Q_o$  even in low canopy layers, would result in high early morning conductance and mute stomatal response to a further in-

crease in light (Fig. 2C; Roden and Pearcy 1993; Allen and Pearcy 2000). However, enhanced stomatal sensitivity to D at night keeps water potential high to sustain growth (Landsberg 1986) at a time that low conductance has no impact on carbon uptake.

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#### References

- Allen MT, Pearcy RW (2000) Stomatal behavior and photosynthetic performance under dynamic light regimes in a seasonally dry tropical rain forest. Oecologia 122:470–478
- Aylmore LAG (1993) CAT studies of water movement. Adv Agron 49:1–54
- Becker P (1998) Limitations of a heat pulse velocity system at low sap flow: implications for measurements at night and in shaded trees. Tree Physiol 18:177–184
- Benyon RG (1999) Nighttime water use in an irrigated *Eucalyptus* grandis plantation. Tree Physiol 19:853–859
- Boyer JS (1989) Water potential and plant metabolism: comments on Dr. P.J. Kramer's article, 'Changing concepts regarding plant water relations', volume 11, pp 565–568, and Dr. J.C. Passioura's Response, pp. 569–571. Plant Cell Environ 12: 213–216
- Bréda N, Cochard H, Dreyer E, Granier A.(1993) Water transfer in a mature oak stand (*Quercus petraea*): seasonal evolution and effects of a severe drought. Can J For Res 23:1136–1143
- Brown CA, Montz GN (1986) Baldcypress. Claitor's, Baton Rouge
- Brown S (1981) A comparison of the structure, primary productivity, and transpiration of cypress ecosystems in Florida. Ecol Monogr 51:403–427
- Bunce JA (1998) Effects of environment during growth on the sensitivity of leaf conductance to changes in humidity. Global Change Biol 4:269–274
- Campbell GS, Norman JM (1998) Environmental biophysics. Springer, Berlin Heidelberg New York
- Carbon BA, Bartle GA, Murray AM, Macpherson DK (1980) The distribution of root length, and the limits to flow of soil water to roots in dry sclerophyll forest. For Sci 26:656–664
- Cienciala E, Lindroth A (1995) Gas-exchange and sap flow measurements of *Salix viminalis* trees in short rotation forest. II. Diurnal and seasonal variation of stomatal response and water use efficiency. Trees 9:295–301
- Cochard H, Breda N, Granier A (1996) Whole tree hydraulic conductance and water loss regulation in *Quercus* during drought: evidence for stomatal control of embolism? Ann Sci For 53:197–206
- Dai Z, Edwards GE, Ku MSB (1992) Control of photosynthesis and stomatal conductance in *Ricinus communis* by leaf to air vapor pressure deficit. Plant Physiol 99:1426–1434
- Dang Q-L, Margolis HA, Coyea MR, Mikailou SY, Collatz GJ (1997) Regulation of branch-level gas exchange of boreal trees: roles of shoot water potential and vapor pressure difference. Tree Physiol 17:521–535
- Doley D (1967) Water relations of *Eucalyptus marginata* Sm. under natural conditions. J Ecol 55:597–614
- Ewers BE, Oren R (2000) Analysis of assumptions and errors in the calculation of stomatal conductance from sap flux measurements. Tree Physiol 20:579–589
- Friend AD (1993) The predictions and physiological significance of tree height. In: Solomon AM, Shugart HH (eds) Vegetation dynamics & global change. Chapman & Hall, New York, pp 101–115

- Gardingen PR van, Grace J, Jeffree CE (1991) Abrasive damage by wind to the needle surfaces of *Picea sitchensis* (Bong.) Carr. and *Pinus sylvestris* L. Plant Cell Environ 14:185–193
- Grace J (1989) Measurement of wind speed near vegetation. In: Pearcy RW, Ehleringer J, Mooney HA, Rundel PW (eds) Plant physiological ecology: field methods and instrumentation. Chapman & Hall, London, pp 57–74
- Granier A (1987) Evaluation of transpiration in Douglas-fir stand by means of sap flow measurements. Tree Physiol 3:309– 320
- Granier A, Bréda N (1996) Modeling canopy conductance and stand transpiration of an oak forest from sap flow measurements. Ann Sci For 53:537–546
- Granier A, Loustau D (1994) Measuring and modeling the transpiration of a maritime pine canopy from sap flow data. Agric For Meteorol 51:309–319
- Granier A, Biron P, Bréda N, Pontailler J-Y, Saugier B (1996) Transpiration of trees and forest stands: short and long-term monitoring using sapflow methods. Global Change Biol 2:265–274
- Green SR, McNaughton KG, Clothier BE (1989) Observations of nighttime water use in kiwifruit vines and apple trees. Agric For Meteorol 48:251–261
- Hadley JH, Smith WK (1990) Influence of leaf surface wax and leaf area to water content ratio on cuticular transpiration in western conifers, U.S.A. Can J For Res 20:1306–1311
- Hinckley TM, Lassoie JP, Running SW (1978) Temporal and spatial variations in water status of forest trees. For Sci Monogr 20
- Hogg EH, Hurdle PA (1997) Sap flow in trembling aspen: implications for stomatal responses to vapor pressure deficit. Tree Physiol 17:501–509
- Iritz Z, Lindroth A (1994) Night-time evaporation from a shortrotation willow stand. J Hydrol 157:235–245
- Jarvis PG (1976) The interpretation of the variations in leaf water potential and stomatal conductance found in canopies in the field. Phil Trans R Soc Lond B 273:593–610
- Jones HG (1992) Plants and microclimate: a quantitative approach to environmental plant physiology. Cambridge University Press, Cambridge, UK
- Körner C (1993) Leaf diffusive conductance in the major vegetation types of the globe.Ecol Stud 100:463–490
- Köstner B, Schulze E-D, Kelliher FM, Hollinger DY, Byers JN, Hunt JE, McSeveny TM, Meserth R, Weir PL (1992) Transpiration and canopy conductance in a pristine broad-leaved forest of *Nothofagus*: an analysis of sapflow and eddy correlation measurements. Oecologia 91:350–359
- Kramer PJ (1983) Water relations of plants. Academic Press, New York
- Kramer PJ (1988) Changing concepts regarding plant water relations. Plant Cell Environ 11:565–568
- Landsberg JJ (1986) Physiological ecology of forest production. Academic Press, London
- Leuning R (1995) A critical appraisal of a combined stomatalphotosynthesis model for  $C_3$  plants. Plant Cell Environ 18: 339–355
- Lohammar T, Larsson S, Linder S, Falk SO (1980) FAST simulation models of gaseous exchange in Scots pine. Ecol Bull (Stockholm) 32:505–523
- Lu P, Biron P, Granier A, Cochard H (1996) Water relations of Norway spruce (*Picea abies* (L) Karst.) under soil drought in the Vosges mountains: whole-tree hydraulic conductance, xylem embolism and water loss regulation. Ann Sci For 53:113–121
- Martin TA, Brown KJ, Cermák J, Ceulemans R, Kucera J, Meinzer FC, Rombold JS, Sprugel DG, Hinckley TM (1997) Crown conductance and tree and stand transpiration in a second growth *Abies amabilis* forest. Can J For Res 27: 797–808
- Massman WJ, Kaufmann MR (1991) Stomatal response to certain environmental factors: a comparison of models for subalpine trees in the Rocky Mountains. Agric For Meteorol 54:155–167

- McCaughey JH, Iacobelli A (1994) Modelling stomatal conductance in a northern deciduous forest, Chalk River, Ontario. Can J For Res 24:904–910
- McCully ME, Huang CX, Ling LEC (1998) Daily embolism and refilling of xylem vessels in the roots of field-grown maize. New Phytol 138:327–342
- McNaughton KG, Jarvis PG (1991) Effects of spatial scale on stomatal control of transpiration. Agric For Meteorol 54:279–301
- Megonigal JP, Conner WH, Kroeger S, Sharitz RR (1997) Aboveground production in southeastern floodplain forests: a test of the subsidy-stress hypothesis. Ecology 78:370–384
- Meinzer FC (1993) Stomatal control of transpiration. Trends Ecol Evol 8:289–293
- Meinzer FC. Hinckley TM, Ceulemans R (1997) Apparent responses of stomata to transpiration and humidity in a hybrid poplar canopy. Plant Cell Environ 20:1301–1308
- Monteith JL (1995) A reinterpretation of stomatal response to humidity. Plant Cell Environ 18:357–364
- Monteith JL, Unsworth MH (1990) Principles of environmental physics. Arnold, London
- Mott KA, Parkhurst DF (1991) Stomatal response to humidity in air and helox. Plant Cell Environ 14:509–515
- Oren R, Phillips N, Katul G, Ewers BE, Pataki DE (1998a) Scaling xylem sap flux and soil water balance and calculating variance: a method for partitioning water flux in forests. Ann Sci For 55:191–216
- Oren R, Ewers BE, Todd P, Phillips N, Katul G (1998b) Water balance delineates the soil layer in which moisture affects canopy conductance. Ecol Appl 8:990–1002
- Oren R, Phillips N, Ewers BE, Pataki DE, Megonigal JP (1999a) Responses of sap flux-scaled transpiration to light, vapor pressure deficit, and leaf area reduction in a flooded *Taxodium distichum* L. forest. Tree Physiol 19:337–347
- Oren R, Sperry JS, Katul GG, Ewers BE, Pataki DE, Phillips N, Schäfer KVR (1999b) Survey and synthesis of intra- and interspecific responses of canopy stomatal conductance to vapour pressure deficit. Plant Cell Environ 22:1515–1526
- Parker GG (1995) Structure and microclimate of forest canopies. In: Lowman MD, Nadkarni NM (eds) Forest canopies. Academic Press, San Diego, pp 73–106
  Pataki DE, Oren R, Phillips N (1998) Responses of sap flux and
- Pataki DE, Oren R, Phillips N (1998) Responses of sap flux and stomatal conductance of *Pinus taeda* L. to stepwise reductions in leaf area. J Exp Bot 49:871–878
- Pearcy RW, Schulze E-D, Zimmermann R (1989) Measurement of transpiration and leaf conductance. In: Pearcy RW, Ehleringer J, Mooney HA, Rundel PW (eds) Plant physiological ecology. Chapman & Hall, London, pp 137–160
- Pearcy RW, Chazdon RL, Gross LJ, Mott KA (1994) Photosynthetic utilization of sunflecks: a temporally patchy resource on a time scale of seconds to minutes. In: Caldwell MM, Pearcy RW (eds) Exploitation of environmental heterogeneity by plants. Academic Press, San Diego, pp 175–208
- Pezeshki SR, Anderson PH (1997) Responses of three bottomland species with different flood tolerance capabilities to various flooding regimes. Wetlands Ecol Manage 4:245–256
- Pezeshki SR, DeLaune RD, Choi HS (1995) Gas exchange and growth of bald cypress seedlings from selected U.S. Gulf Coast populations: responses to elevated salinities. Can J For Res 25:1409–1415
- Phillips N, Oren R (1998) A comparison of daily representations of canopy conductance based on two conditional time-averaging methods and the dependence of daily conductance on environmental factors. Ann Sci For 55:217–235
- Phillips N, Oren R, Zimmermann R (1996) Radial patterns of xylem sap flow in non-, diffuse- and ring-porous tree species. Plant Cell Environ 19:983–990
- Putz FE, Sharitz RR (1991) Hurricane damage to old-growth forest in Congaree swamp National Monument, South Carolina, U.S.A. Can J For Res 21:1765–1770
- Roden JS, Pearcy RW (1993) Photosynthetic gas exchange response of poplars to steady-state and dynamic light environments. Oecologia 93:208–214

- Saliendra NZ, Sperry JS, Comstock JP (1995) Influence of leaf water status on stomatal response to humidity, hydraulic conductance, and soil drought in *Betula occidentalis*. Planta 196:357–366
- Sandford AP, Jarvis PG (1986) Stomatal responses to humidity in selected conifers. Tree Physiol 2:89–103
- Schäfer KVR, Oren R, Tenhunen JD (2000) The effect of tree height on crown-level stomatal conductance. Plant Cell Environ 23:365–375
- Schulze E-D, Cermák J, Matyssek R, Penka M, Zimmermann R, Vasícek F, Gries W, Kucera J (1985) Canopy transpiration and water fluxes in the xylem of the trunk of *Larix* and *Picea* trees – a comparison of xylem flow, porometer and cuvette measurements. Oecologia 66:475–483
- Schulze E-D, Kelliher FM, Körner C, Lloyd J, Leuning R (1994) Relationship among maximum stomatal conductance, ecosystem surface conductance, carbon assimilation rate, and nitrogen nutrition: a global ecology scaling exercise. Ann Rev Ecol Syst 25:629–660
- Sena Gomes AR, Kozlowski TT (1980) Growth responses and adaptations of *Fraxinus pensylvanica* seedlings to flooding. Plant Physiol 66:267–271
- Sperry JS (1995) Limitations on stem water transport and their consequences. In: Gardner B (ed) Plant stems. Academic Press, San Diego, pp 105–124

- Sperry JS, Alder NN, Eastlack SE (1993) The effect of reduced hydraulic conductance on stomatal conductance and xylem cavitation. J Exp Bot 44:1075–1082
- Sperry JS, Adler FR, Campbell GS, Comstock JC (1998) Limitation of plant water use by rhizosphere and xylem conductance: results from a model. Plant Cell Environ 21:347–359
- Steinberg SL, Bavel CHM van, McFarland MJ (1990) Improved sap flow gauge for woody and herbaceous plants. Agron J 83: 851–854
- Sullivan NH, Bolstad PV, Vose JM (1996) Estimates of net photosynthetic parameters for twelve tree species in mature forests of the southern Appalachians. Tree Physiol 16:397–406
- Tyree MT, Sperry JS (1988) Do woody plants operate near the point of catastrophic xylem dysfunction caused by dynamic water stress? Answers from a model. Plant Physiol 88:574–580
- Tyree MT, Cochard H, Cruiziat P, Sinclair B, Ameglio T (1993) Drought induced leaf shedding in walnut: evidence for vulnerability segmentation. Plant Cell Environ 16:879–882
- Whitehead D (1998) Regulation of stomatal conductance and transpiration in forest canopies. Tree Physiol 18:633–644
- Whitehead D, Jarvis PG (1981) Coniferous forests and plantations. In: Kozlowski TT (ed) Water deficits and plant growth. Academic Press, Orlando, pp 49–149
- Wurman J, Winslow J (1998) Intense sub-kilometer-scale boundary layer rolls observed in Hurricane Fran. Science 280: 555–557